06/27/2006 16:39 703-385-9719 KILYK & BOWERSOX PAGE 25

Application No. 10/672,937 Amendment dated June 27, 2006

Response to Office Action dated December 27, 2006

Amendments to the Drawings:

The attached two sheets of drawings include changes to Figs. 5 and 6, and replace the original sheets including Figs. 5 and 6. Figs. 5 and 6 have been amended to delete the text labels "Table 1" and "Table 2," respectively.

Attachment:

Replacement Sheet

Annotated Sheet Showing Changes

Application No. 10/672,937 Amendment dated June 27, 2006 Response to Office Action dated December 27, 2006

REMARKS

INTRODUCTION

This Amendment is in response to the Office Action dated December 27, 2005. Applicants sincerely thank Examiner Zhou for the courtesies extended to Applicants' representative, Scott Balderston, during the personal interview conducted June 15, 2006. Applicants have amended the application in light of the Office Action to address all objections to the specification and drawings, and all rejections of the claims. Favorable reconsideration is respectfully requested.

OBJECTIONS TO THE SPECIFICATION AND DRAWINGS

The Office Action objected to the title of the invention. Applicants have amended the title in response to that objection. The Office Action objected to the reference to "AB1" format for certain computer data noted, for instance, at pages 21 and 30 of the specification. Applicants respectfully submit that the file extension or file identifier "AB1" is in fact a type of file format or identifier known in the art. Applicants provide with this Amendment an Information Disclosure Statement submitting an excerpt from a document entitled "ABI Prism® 3000 Genetic Analyzer" making a notation of that file format at page 5-9.

The Office Action objected to the Internet hyperlink "www.phrap.org" referenced at page 3, line 14, of the specification. Applicants have truncated the reference to "phrap.org" so that reference does not represent a full embedded hyperlink. The Office Action objected to the drawings and specification because the data shown in Figs. 5 and 6 were labeled as "Table 1" and "Table 2"

Page 20 of 23

Application No. 10/672,937 Amendment dated June 27, 2006 Response to Office Action dated December 27, 2006

respectively, but shown as figures in the drawings. Applicants have deleted the labels "Table 1" and "Table 2" in Figs. 5 and 6 and conformed the specification to refer to Figs. 5 and 6, accordingly.

Applicants have therefore considered and duly responded to all of the objections to the specification and drawings, which are therefore overcome. Reconsideration and withdrawal of the objections are respectfully requested.

REJECTIONS OF CLAIMS UNDER 35 U.S.C. §112, SECOND PARAGRAPH

The Office Action rejected claims 1-18 and 43 under 35 U.S.C. §112, second paragraph for indefiniteness due to the term "reportable ranges" in claims 1 and 43. Applicants have amended claims 1 and 43 to recite "wherein the reportable ranges comprise base sequence ranges related to the at least one sample, usable for comparison matching". As discussed in the specification, for instance, at page 19, lines 10-22 and at page 26, line 11 – page 27, line 2, those "reportable ranges" can, in various embodiments and generally speaking, represent DNA, mtDNA, or other base sequence ranges that have been determined to be sufficiently unambiguous or reliable to use for comparison matching purposes. Claims 1-18 and 43 are therefore adequately clear, and it is respectfully submitted that the rejection is therefore overcome.

Applicants understand the Office Action to have rejected all dependent claims which recite the term "major and minor bases," under 35 U.S.C. §112, second paragraph, as being indefinite. Claims 2 and 4-12 recite that term. Applicants respectfully submit that the term "major and minor bases" would have been adequately clear to a person skilled in the art at the time the application was filed, so that this rejection is overcome. Applicants submit that "major and minor

Page 21 of 23

06/27/2006 16:39 703-385-9719 KILYK & BOWERSOX PAGE 28

Application No. 10/672,937 Amendment dated June 27, 2006

Response to Office Action dated December 27, 2006

bases" would be understood by persons skilled in the art to refer to bases that are associated with

major peaks and minor peaks of an electropherogram generated by base-calling or sequencing

operations. Applicants provide with this Amendment an Information Disclosure Statement

submitting a document entitled "Comparison of PowerPlexTM 16, Plower PlexTM 1.1/2.1, and ABI

Ampf/STRTM Profiler PlusTM/COfilerTM for Forensic Use". That document refers, for example, at

page 42:241, to the association of alleles with major and minor peaks in electropherograms

generated by ABI PrismTM hardware, consistent with this connotation. Various peaks generated by

such operations are, for example, illustrated in Applicants' Fig. 1, and described, for instance, in the

specification at page 16, lines 16-22. Applicants respectfully submit that the term "major and minor

bases" is adequately clear, and that the rejection is overcome.

The Office Action rejected claim 18 under 35 U.S.C. §112, second paragraph, as being

indefinite due to the language "wherein the sequence information corresponds to mitochondrial

sequence information." Applicants have amended this language to recite that the sequence

information "comprises mitochondrial sequence information." All rejections of claims 1-18 and 43,

under 35 U.S.C. §112, second paragraph, are thereby duly addressed and overcome.

Reconsideration and withdrawal of the rejection are respectfully requested.

CONCLUSION

In view of the foregoing remarks, Applicants respectfully request favorable reconsideration

of the present application and a timely allowance of the pending claims.

Page 22 of 23

Application No. 10/672,937 Amendment dated June 27, 2006 Response to Office Action dated December 27, 2006

Should the Examiner deem that any further action by Applicants or Applicants' undersigned representative is desirable and/or necessary, the Examiner is invited to telephone the undersigned at the number set forth below.

If there are any other fees due in connection with the filing of this response, please charge the fees to deposit Account No. 50-0925. If a fee is required for an extension of time under 37 C.F.R. § 1.136 not accounted for above, such extension is requested and should also be charged to said Deposit Account.

Respectfully submitted,

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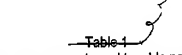
Enclosures: Corrected Fogs. 5 and 6

Annotated Version Showing Corrections to Figs. 5 and 6

ANNOTATED SHEET

AL CHONDRIAL DNA AUTOSCORING SYSTEM Stockwell et al.

Alty Docket: ABIOS.042A Appl. No.: Unknown



System components and tunable parameters

Program Punction performed
mark_substitution_heteroplasmy.pl Mark ambiguous base calls extract_SE_consensus.pl Extract a consensus sequence from phrap polyphred run, replacing low quali bases with '?', and polyphred rank bases with 'N'. Calculates overall, forward strand, ar reverse strand coverage for all bases the consensus sequence Count_hv1_deletes.pl Compute start and end positions of HV and HV2 regions.
extract_SE_consensus.pl Extract a consensus sequence from phrap polyphred run, replacing low quality bases with '?', and polyphred rank bases with 'N'. Compute_coverage.pl Calculates overall, forward strand, ar reverse strand coverage for all bases the consensus sequence Count_hv1_deletes.pl Compute start and end positions of HV and HV2 regions.
bases with '?', and polyphred rank bases with 'N'. Compute_coverage.pl Calculates overall, forward strand, ar reverse strand coverage for all bases the consensus sequence Count_hv1_deletes.pl Count deletions in HV1 relative to the rCRS border_index.pl Compute start and end positions of HV and HV2 regions.
bases with 'N'. Calculates overall, forward strand, ar reverse strand coverage for all bases the consensus sequence Count_hv1_deletes.pl Count_deletions in HV1 relative to the rCRS border_index.pl Compute start and end positions of HV and HV2 regions.
compute_coverage.pl Calculates overall, forward strand, ar reverse strand coverage for all bases the consensus sequence count_hv1_deletes.pl Count_deletions in HV1 relative to the rCRS border_index.pl Compute start and end positions of HV and HV2 regions.
reverse strand coverage for all bases the consensus sequence count_hv1_deletes.pl
the consensus sequence count_hv1_deletes.pl Count_deletions in HV1 relative to the rCRS border_index.pl Compute start and end positions of HV and HV2 regions.
count_hv1_deletes.pl Count_deletions in HV1 relative to the rCRS border_index.pl Compute start and end positions of HV and HV2 regions.
border_index.pl Compute start and end positions of HV and HV2 regions.
and HV2 regions.
and HV2 regions.
fix mitotype reporting range of Output a mtDNA profile based on inp
variants list and reportable range
find_bad_traces_from_blast_report.pl Reject reads that do not alignment alignment.
appropriately to the rCRS
determineReadTypes.pl Adds template name, template type, at primer type to phred output files.
seq2delta_vs Align intDNA profile to rCRS and report variants according to nomenclature.
calculate_coverage_mitotype.pl Mask mtDNA consensus sequence base
on required coverage and calculate
coverage
count_hv1_inserts.pl Count insertions in HV1 relative to the
rCRS
generate_hv_mask_fasta_files.pl
of the mtDNA consensus sequence based on the computed HV1 and HV
regions.
flip_fasta.pl Invert an X masked sequence output
cross_match

ANNOTATED SHEET

M HONDRIAL DNA AUTUSCURING 3131 Em.
Stockwell et al.

Appl. No.: Unknown Atty Docket: ABIOS.042A

Table 2

	Assessment 1	<u> Assessment 2</u>
Total True Positives: N _{sutomated} = N _{mmunl}	16	10
Total True Negatives: Apulimated = Amanual	77,358	110,354
Total False Positives: Nautomated = Amenual	95	807
Total False Negatives: Antipmated = Nmanual	6ª	14
Total Incorrect	0	0
Sensitivity: TP/(TP+FN)	72.73%	41.67%
Specificity: TN/(TN+FP)	99.88%	99.27%
Positive Predictive Value: TP/(TP+FP)	14.41%	1.22%
Negative Predictive Value: TN/(TN+FN)	99.99%	99.99%

^aPredominate base correctly called

Differences take into account the consensus of the two analysts